

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE
 REQUEST FOR FILING NATIONAL PHASE OF
 PCT APPLICATION UNDER 35 U.S.C. 371 AND 37 CFR 1.494 OR 1.495

09/857766

To: Hon. Commissioner of Patents
 Washington, D.C. 20231



00909

TRANSMITTAL LETTER TO THE UNITED STATES
 DESIGNATED/ELECTED OFFICE (DO/EO/US)

Atty Dkt: P 281380 /SMC-60328/UST
 M# /Client Ref.

From: Pillsbury Winthrop LLP, IP Group:

Date: June 11, 2001

This is a **REQUEST** for **FILING** a PCT/USA National Phase Application based on:

1. International Application PCT/GB99/03807 1 country code	2. International Filing Date 16 November 1999 Day MONTH Year	3. Earliest Priority Date Claimed 12 December 1998 Day MONTH Year (use item 2 if no earlier priority)
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4. Measured from the earliest priority date in item 3, this PCT/USA National Phase Application Request is being filed within:

(a) ☐ 20 months from above item 3 date (b) ☒ 30 months from above item 3 date,

(c) Therefore, the due date (unextendable) is June 12, 2001

Title of Invention COMPOSITION AND PROCESS FOR THE EXTRACTION OF METALS

Inventor(s) SUGARMAN, Alan David

Applicant herewith submits the following under 35 U.S.C. 371 to effect filing:

☒ Please immediately start national examination procedures (35 U.S.C. 371 (f)).

☒ A copy of the International Application as filed (35 U.S.C. 371(c)(2)) is transmitted herewith (file if in English but, if in foreign language, file only if not transmitted to PTO by the International Bureau) including:

a. ☒ Request;

b. ☒ Abstract;

c. 17 pgs. Spec. and Claims;

d. _____ sheet(s) Drawing which are ☐ informal ☐ formal of size ☐ A4 ☐ 11"

9. ☒ A copy of the International Application has been transmitted by the International Bureau.

10. A translation of the International Application into English (35 U.S.C. 371(c)(2))

a. ☐ is transmitted herewith including: (1) ☐ Request; (2) ☐ Abstract;

(3) _____ pgs. Spec. and Claims;

(4) _____ sheet(s) Drawing which are:

☐ informal ☐ formal of size ☐ A4 ☐ 11"

b. ☐ is not required, as the application was filed in English.

c. ☐ is not herewith, but will be filed when required by the forthcoming PTO Missing Requirements Notice per Rule 494(c) if box 4(a) is X'd or Rule 495(c) if box 4(b) is X'd.

d. ☐ Translation verification attached (not required now).

09/057766
11 JUN 2001

11. ☒ Please see the attached Preliminary Amendment
12. ☐ Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3)), i.e., before 18th month from first priority date above in item 3, are transmitted herewith (file only if in English) including:
13. ☒ PCT Article 19 claim amendments (if any) have been transmitted by the International Bureau
14. ☐ Translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)), i.e., of claim amendments made before 18th month, is attached (required by 20th month from the date in item 3 if box 4(a) above is X'd, or 30th month if box 4(b) is X'd, or else amendments will be considered canceled).

15. **A declaration of the inventor** (35 U.S.C. 371(c)(4))

- a. ☒ is submitted herewith ☒ Original ☐ Facsimile/Copy
- b. ☐ is not herewith, but will be filed when required by the forthcoming PTO Missing Requirements Notice per Rule 494(c) if box 4(a) is X'd or Rule 495(c) if box 4(b) is X'd.

16. **An International Search Report (ISR):**

- a. Was prepared by ☒ European Patent Office ☐ Japanese Patent Office ☐ Other
- b. ☒ has been transmitted by the international Bureau to PTO.
- c. ☒ copy herewith (2 pg(s).) ☒ plus Annex of family members (2 pg(s).).

17. **International Preliminary Examination Report (IPER):**

- a. ☒ has been transmitted (if this letter is filed after 28 months from date in item 3) in English by the International Bureau with Annexes (if any) in original language.
- b. ☒ copy herewith in English.
- c.1 ☐ IPER Annex(es) in original language ("Annexes" are amendments made to claims/spec/drawings during Examination) including attached amended:
- c.2 ☐ Specification/claim pages # _____ claims # _____ Dwg Sheets # _____
- d. ☐ Translation of Annex(es) to IPER (required by 30th month due date, or else annexed amendments will be considered canceled).

18. **Information Disclosure Statement** including:

- a. ☒ Attached Form PTO-1449 listing documents
- b. ☒ Attached copies of documents listed on Form PTO-1449
- c. ☒ A concise explanation of relevance of ISR references is given in the ISR.

19. ☒ **Assignment** document and Cover Sheet for recording are attached. Please mail the recorded assignment document back to the person whose signature, name and address appear at the end of this letter.20. ☐ Copy of Power to IA agent.21. ☐ **Drawings** (complete only if 8d or 10a(4) not completed): ___ sheet(s) per set: ☐ 1 set informal; ☐ Formal of size ☐ A4 ☐ 11"22. Small Entity Status ☒ is **Not** claimed ☐ is claimed (pre-filing confirmation required)

22(a) _____ (No.) Small Entity Statement(s) enclosed (since 9/8/00 Small Entity Statements(s) not essential to make claim)

23. **Priority** is hereby claimed under 35 U.S.C. 119/365 based on the priority claim and the certified copy, both filed in the International Application during the international stage based on the filing in (country) **GREAT BRITAIN** of:

	Application No.	Filing Date		Application No.	Filing Date
(1)	9827288.3	Dec. 12, 1998	(2)	_____	_____
(3)	_____	_____	(4)	_____	_____
(5)	_____	_____	(6)	_____	_____

- a. ☒ See Form PCT/IB/304 sent to US/DO with copy of priority documents. If copy has not been received, please proceed promptly to obtain same from the IB.
- b. ☐ Copy of Form PCT/IB/304 attached.

24. Attached:

25 Per Item 17.c2, **cancel original** pages #____, claims #____, Drawing Sheets #26. **Calculation of the U.S. National Fee (35 U.S.C. 371 (c)(1)) and other fees is as follows:**Based on amended claim(s) per above item(s) ☐ 12, ☐ 14, ☐ 17, ☐ 25 (hilitte)

Total Effective Claims	23	minus 20 =	3	x \$18/\$9	=	\$54	966/967
Independent Claims	2	minus 3 =	0	x \$80/\$40	=	\$0	964/965
If any proper (ignore improper) Multiple Dependent claim is present,				add \$270/\$135	=	+270	968/969

BASIC NATIONAL FEE (37 CFR 1.492(a)(1)-(4)): →→ **BASIC FEE REQUIRED, NOW** →→→→A. If country code letters in item 1 are **not** "US", "BR", "BB", "TT", "MX", "IL", "NZ", "IN" or "ZA"

See item 16 re:

1. Search Report was <u>not prepared</u> by EPO or JPO -----	add \$1000/\$500		960/961
2. Search Report was prepared by EPO or JPO -----	add \$860/\$430	+860	970/971

SKIP B, C, D AND E UNLESS country code letters in item 1 are "US", "BR", "BB", "TT", "MX", "IL", "NZ", "IN" or "ZA"

→ <input type="checkbox"/> B. If USPTO did not issue <u>both</u> International Search Report (ISR) <u>and</u> (if box 4(b) above is X'd) the International Examination Report (IPER), -----	add \$1000/\$500	+0	960/961
→ <input type="checkbox"/> C. If USPTO issued ISR but not IPER (or box 4(a) above is X'd), -----	add \$710/\$355	+0	958/959
→ <input type="checkbox"/> D. If USPTO issued IPER but IPER Sec. V boxes <u>not all</u> 3 YES, -----	add \$690/\$345	+0	956/957
→ <input type="checkbox"/> E. If international preliminary examination fee was paid to USPTO and Rules 492(a)(4) and 496(b) <u>satisfied</u> (IPER Sec. V <u>all</u> 3 boxes YES for <u>all</u> claims), -----	add \$100/\$50	+0	962/963
SUBTOTAL =		\$1184	

28. If Assignment box 19 above is X'd, add Assignment Recording fee of ---\$40	+40	(581)
29. Attached is a check to cover the -----	TOTAL FEES	\$1224

Our Deposit Account No. 03-3975

Our Order No. 70662 281380

C#

M#



CHARGE STATEMENT: The Commissioner is hereby authorized to charge any fee specifically authorized hereafter, or any missing or insufficient fee(s) filed, or asserted to be filed, or which should have been filed herewith or concerning any paper filed hereafter, and which may be required under Rules 16-18 and 492 (missing or insufficient fee only) now or hereafter relative to this application and the resulting Official document under Rule 20, or credit any overpayment, to our Account/Order Nos. shown above for which purpose a duplicate copy of this sheet is attached.

This CHARGE STATEMENT does not authorize charge of the issue fee until/unless an issue fee transmittal form is filedPillsbury Winthrop LLP
Intellectual Property GroupBy Atty: Paul N. Kokulis Reg. No. 16773Sig: [Signature] Fax: (202) 822-0944
Tel: (202) 861-3503

Atty/Sec: PNK/mhn

NOTE: File in duplicate with 2 postcard receipts (PAT-103) & attachments.

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59/541/0077 11 JUN 2001**IN THE UNITED STATES PATENT AND TRADEMARK OFFICE**

In re PATENT APPLICATION OF

Inventor(s): SUGARMAN, Alan David

Filed: Herewith

Title: COMPOSITION AND PROCESS FOR THE EXTRACTION OF METALS

June 11, 2001

PRELIMINARY AMENDMENTHon. Commissioner of Patents
Washington, D.C. 20231

Sir:

Please amend this application as follows:

IN THE SPECIFICATION:

At the top of the first page, just under the title, insert

☒ --This application is the National Phase of International Application
PCT/GB99/03807 filed November 16, 1999 which designated the U.S.
and that International Application

☒ was ☐ was not published under PCT Article 21(2) in English.--

Respectfully submitted,

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COMPOSITION AND PROCESS FOR THE EXTRACTION OF METALS

The present invention concerns a solvent extraction composition, a solvent extraction process and especially a process for the extraction of metals, particularly copper, from aqueous solutions, especially solutions obtained by leaching ores.

It is known to extract metals, especially copper, from aqueous solutions containing the metal in the form of, for example, a salt, by contacting the aqueous solution with a solution of a solvent extractant in a water immiscible organic solvent and then separating the solvent phase loaded with metal, i.e. containing at least a part of the metal in the form of a complex. The metal can then be recovered by stripping with a solution of lower pH followed for example, by electrowinning. Most commonly, the aqueous metal-containing solutions for extraction are the result of the acid leaching of ores. However it is known that some metals, especially copper, can be leached from certain ores with ammoniacal solutions. This has the advantage that solutions containing especially high concentrations of copper are derived and that there is little contamination of the solution with iron.

Solvent extractants which have found favour in recent years particularly for the recovery of copper from aqueous solutions include oxime reagents, especially o-hydroxyaryloximes and o-hydroxyarylketo oximes. Whilst such reagents have been found to work well in the recovery of copper from solutions, one problem which has been encountered in the application of such reagents is that the oxime and keto xime reagents can strongly bind metals to the extent that the efficiency of metal transfer from leach solution to strip solution can be impaired. In order to overcome such problems, modifiers have been used to effect the binding efficiency of the extractants. Typical modifiers are disclosed in WO96/25525, and in particular a class of highly branched ester modifiers are disclosed in EP-A-0202833. However, as solvent extraction processes are increasingly employed in more diverse situations, there is still a need to identify further modifiers.

According to a first aspect of the present invention, there is provided a solvent extraction composition comprising one or more orthohydroxyaryldoximes or orthohydroxyarylketo oximes, and one or more esters substituted with a hydroxy group. The compositions preferably also comprise a water immiscible organic solvent.

The orthohydroxyaryldoxime or orthohydroxyarylketo xime compounds employed in the present invention are substantially water insoluble and preferably have the formula:



Formula (1)

wherein

R¹ is hydrogen or an optionally substituted hydrocarbonyl group

R² is an optionally substituted ortho-hydroxyaryl group,
and salts thereof.

Whilst the invention is described herein with reference to a compound of Formula (1), it is understood that it relates to said compound in any possible tautomeric forms, and also the complexes formed between orthohydroxyarylaloximes or orthohydroxyarylketoimes and metals, particularly copper.

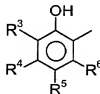
Optionally substituted hydrocarbonyl groups which may be represented by R¹ preferably comprise optionally substituted alkyl and aryl groups including combinations of these, such as optionally substituted aralkyl and alkaryl groups.

Examples of optionally substituted alkyl groups which may be represented by R¹ include groups in which the alkyl moieties can contain from 1 to 20, especially from 1 to 4, carbon atoms. A preferred orthohydroxyarylketoime is one in which R¹ is alkyl, preferably containing up to 20, and especially up to 10, and more preferably up to 3 saturated aliphatic carbon atoms, and most preferably R¹ is a methyl group.

Examples of optionally substituted aryl groups which may be represented by R¹ include optionally substituted phenyl groups. When R¹ is an aryl group, it is preferably an unsubstituted phenyl group.

Most preferably R¹ represents a hydrogen atom.

Optionally substituted ortho-hydroxyaryl groups which may be represented by R² include optionally substituted phenols. Examples of optionally substituted phenols which may be represented by R² include those of formula:



wherein R³ to R⁶ each independently represent H or a C₁ to C₂₂, preferably a C₇ to C₁₅, linear or branched alkyl group. Particularly preferably only R⁵ represents a C₁₋₂₂ alkyl group, most preferably a C₇ to C₁₅ alkyl group, with R³, R⁴ and R⁶ representing H.

When R¹ or R² is substituted, the substituent(s) should be such as not to affect adversely the ability of the orthohydroxyarylaloxime or orthohydroxyarylketoime to complex with metals, especially copper. Suitable substituents include halogen, nitro, cyano, hydrocarbonyl, such as C₁₋₂₀-alkyl, especially C₁₋₁₀-alkyl; hydrocarbonyloxy, such as C₁₋₂₀-alkoxy, especially C₁₋₁₀-alkoxy; hydrocarbonyloxycarbonyl, such as C₁₋₂₀-alkoxycarbonyl, especially C₁₋₁₀-alkoxycarbonyl; acyl, such as C₁₋₂₀-alkylcarbonyl and arylcarbonyl, especially C₁₋₁₀-alkylcarbonyl and phenylcarbonyl; and acyloxy, such as C₁₋₂₀-alkylcarbonyloxy and arylcarbonyloxy, especially C₁₋₁₀-alkylcarbonyloxy and

phenylcarbonyloxy. There may be more than one substituent in which case the substituents may be the same or different.

In many embodiments, when an orthohydroxyarylketoxy employed, the orthohydroxyarylketoxy is a 5-(C₈ to C₁₄ alkyl)-2-hydroxyacetophenone oxime, particularly 5-nonyl-2-hydroxyacetophenone oxime.

In many preferred embodiments, when an orthohydroxyarylaldoxy employed, the orthohydroxyarylaldoxy is a 5-(C₈ to C₁₄ alkyl)-2-hydroxybenzaldoxy, particularly 5-nonyl-2-hydroxybenzaldoxy.

The composition may comprise one or more different orthohydroxyarylaldoxys or orthohydroxyarylketoxys or mixtures thereof in which the nature of the substituent groups represented by R¹ and R² differ between component orthohydroxyarylaldoxys or orthohydroxyarylketoxys, especially where the component orthohydroxyarylaldoxys or orthohydroxyarylketoxys are isomeric. Such isomeric mixtures may have better solubility in organic solvents than a single orthohydroxyarylketoxy.

The orthohydroxyarylaldoxys or orthohydroxyarylketoxys are often present in an amount of up to 60% by weight of the composition, commonly no more than 50%, and usually no more than 40 % w/w. Often, the orthohydroxyarylaldoxy or orthohydroxyarylketoxy comprises at least 1% by weight, commonly at least 2.5% by weight and usually at least 5% by weight of composition, and preferably comprises from 7.5 to 20%, such as about 10%, by weight of the composition.

The esters substituted with a hydroxy group employed in the present invention are substantially water insoluble and commonly have the formula:



Formula (2)

wherein one of R⁷ or R⁸ is a substituted hydrocarbyl group which is substituted with at least one hydroxyl group and the other is an optionally substituted hydrocarbyl group. Preferably, the ester is an aliphatic or aromatic-aliphatic ester, and particularly preferably a branched chain aliphatic or aromatic-aliphatic ester.

The esters substituted with a hydroxy group (hereinafter Hydroxy-esters) which may be employed in this invention may be di-esters or polyesters, but are especially mono-esters.

The Hydroxy-esters of the present invention can contain from 5 to 51 carbon atoms, preferably from 7 to 40 carbon atoms, and more preferably from 9 to 25 carbon atoms.

The Hydroxy-esters are preferably highly branched. In the context of the present invention, "highly branched" means that the ratio of the number of methyl carbon atoms

(CH₃ groups) to non-methyl carbon atoms (non CH₃ groups) is higher than 1:5 and preferably higher than 1:3, and is often less than 1.8:1, preferably less than 1.5:1.

Optionally substituted hydrocarbyl groups which may be represented by R⁷ and R⁸ preferably comprise optionally substituted alkyl and aryl groups including combinations of these, such as optionally substituted aralkyl and alkaryl groups.

Examples of optionally substituted alkyl groups which may be represented by R⁷ and R⁸ include groups in which the alkyl moieties can contain from 1 to 25, especially from 2 to 12, carbon atoms. When R⁷ is an optionally substituted alkyl group, it is preferred that it contain up to 12 carbon atoms, more preferably up to 8 carbon atoms, and especially up to 4 carbon atoms. When R⁸ is an optionally substituted alkyl group, it is preferred that it contain at least 2 carbon atoms, especially at least 4 carbon atoms, and especially at least 6 carbon atoms.

Examples of optionally substituted aryl groups which may be represented by R⁷ and R⁸ include optionally substituted phenyl groups. In certain embodiments, only one of R⁷ or R⁸ is an aryl group. It is especially preferred, when only one of R⁷ and R⁸ is an aryl group, that R⁷ is an aryl group. When R⁷ or R⁸ is an aryl group, it is preferably an unsubstituted phenyl group.

The Hydroxy-esters must contain a hydroxy group but optionally may contain other functional groups. Functional groups which may be present include halogen, nitro, cyano, hydrocarbyl, such as C₁₋₂₀-alkyl, especially C₁₋₁₀-alkyl; hydrocarbyloxy, such as C₁₋₂₀-alkoxy, especially C₁₋₁₀-alkoxy; hydrocarbyloxycarbonyl, such as C₁₋₂₀-alkoxycarbonyl, especially C₁₋₁₀-alkoxycarbonyl; acyl, such as C₁₋₂₀-alkylcarbonyl and arylcarbonyl, especially C₁₋₁₀-alkylcarbonyl and phenylcarbonyl; and acyloxy, such as C₁₋₂₀-alkylcarbonyloxy and arylcarbonyloxy, especially C₁₋₁₀-alkylcarbonyloxy and phenylcarbonyloxy. There may be more than one functional group in which case the functional groups may be the same or different.

The hydroxy group that must be present in the Hydroxy-esters of this invention may reside on the portion of the Hydroxy-ester nominally derived from an alcohol (the optionally substituted hydrocarbyl group R⁸) or a carboxylic acid (the optionally substituted hydrocarbyl group R⁷). It is preferred that the hydroxy group reside on the portion of the Hydroxy-ester derived from an alcohol, and that particularly this portion is derivable from a diol. It is especially preferred that the hydroxy group is attached to an aliphatic portion of the Hydroxy-ester.

Where the Hydroxy-ester is a product of the reaction of an alcohol and a hydroxy substituted mono-carboxylic acid, it is preferred that the alcohol is an alkyl alcohol and comprises from 2 to 12 carbon atoms, and the acid is a hydroxy substituted alkyl carboxylic acid and comprises from 2 to 12 carbon atoms. Where the Hydroxy-ester is a product of the reaction of a diol and a mono-carboxylic acid, it is preferred that the diol is an alkyl diol and comprises at least 6 carbon atoms, and the mono-carboxylic acid

comprises from 2 to 12 carbon atoms. Examples of useful esters include 2,2-dimethyl-1,3-propanediol mono-isobutyrate, 2-hydroxymethyl-2-methylpentanol mono-tertpentanoate and nonane-1,3-diol mono-acetate, and particularly 2,2,4-trimethyl-1,3-pentenediol mono-isobutyrate and 2,2,4-trimethyl-1,3-pentenediol monobenzoate.

5 The Hydroxy-esters often comprise up to 30% w/w of the composition, preferably from 0.1 to 20 % w/w, and most preferably from 0.5 to 15% w/w. The weight ratio of Hydroxy-ester to aldoxime or ketoxime is often in the range of from 10:1 to 1:10, commonly from 5:1 to 1:5, and preferably from 1:1 to 1:4.

10 Organic solvents which may be present in the composition include any mobile organic solvent, or mixture of solvents, which is immiscible with water and is inert under the extraction conditions to the other materials present. Examples of suitable solvents include aliphatic, alicyclic and aromatic hydrocarbons and mixtures of any of these as well as chlorinated hydrocarbons such as trichloroethylene, perchloroethylene, trichloroethane and chloroform. Examples of suitable hydrocarbon solvents include
15 ESCAID 110, ESCAID 115, ESCAID 120, ESCAID 200, and ESCAID 300 commercially available from Exxon (ESCAID is a trade mark), SHELLSOL D70 and D80 300 commercially available from Shell (SHELLSOL is a trade mark), and CONOCO 170 commercially available from Conoco (CONOCO is a trade mark). Certain suitable solvents have a low aromatic (<1% w/w) content, for example hydrocarbon solvents such as ESCAID 110 commercially available from Exxon (ESCAID is a trade mark), and
20 ORFOM SX 10 and ORFOM SX11 commercially available from Phillips Petroleum (ORFOM is a trade mark). Preferred solvents are hydrocarbon solvents including high flash point solvents with a high aromatic content such as SOLVESSO 150 commercially available from Exxon (SOLVESSO is a trade mark) and includes solvents which consist
25 essentially of a mixture of trimethylbenzenes such as AROMASOL H, commercially available from Imperial Chemical Industries PLC (AROMASOL is a trade mark). Especially preferred, however, on grounds of low toxicity and wide availability are hydrocarbon solvents of relatively low aromatic content such as kerosene, for example ESCAID 100 which is a petroleum distillate with a total aromatic content of 23%
30 commercially available from Exxon (ESCAID is a trade mark), or ORFOM SX7, commercially available from Phillips Petroleum (ORFOM is a trade mark).

In many embodiments, the composition comprises at least 30%, often at least 45% by weight, preferably from 50 to 95% w/w of water-immiscible hydrocarbon solvent.

35 Certain preferred compositions comprise at least one orthohydroxyaryldoxime or orthohydroxyarylketoxime which may be present in an amount up to 42% w/w, and preferably from 7.5 to 20% w/w, and at least one Hydroxy-ester which may also be present in an amount up to 28%, preferably from 0.5 to 15%, w/w. Compositions comprising at least one orthohydroxyaryldoxime or orthohydroxyarylketoxime which is

present in an amount from 7.5 to 20% w/w and at least one Hydroxy-ester which is present in an amount of from 0.5 to 15% w/w are particularly preferred.

Particularly preferred solvent extraction compositions are those comprising from 7.5 to 20% w/w of 5-(C₈ to C₁₄ alkyl)-2-hydroxyacetophenone oxime or 5-(C₈ to C₁₄ alkyl)-2-hydroxybenzaldoxime, 0.5 to 15% w/w of 2,2,4-trimethyl-1,3-pentanediol mono-isobutryate or benzoate, and from 65 to 92% of water-immiscible hydrocarbon solvent.

Advantageously, it may be preferred to make and supply the composition in the form of a concentrate. In certain embodiments, the concentrate may consist of a mixture of one or more orthohydroxyarylaloximes or orthohydroxyarylketoimes, and one or more Hydroxy-esters (ie no solvent is present). The concentrate may then be diluted by the addition of organic solvents as described herein above to produce compositions in the ranges as described herein above. Where the concentrate contains a solvent, it is preferred that the same solvent is used to dilute the concentrate to the "in use" concentration range. In many embodiments, the concentrate composition comprises up to 30%, often up to 20% by weight, preferably up to 10% w/w of water-immiscible hydrocarbon solvent. Often the concentrate composition comprises greater than 5% w/w of water-immiscible hydrocarbon solvent

If desired, compounds or mixtures of compounds selected from the group consisting of alkylphenols, alcohols, esters, ethers, polyethers, carbonates, ketones, nitriles, amides, carbamates, sulphoxides, and salts of amines and quaternary ammonium compounds may also be employed as additional modifiers in the composition of the invention. Particularly preferred are mixtures comprising a first compound selected from the group consisting of alkylphenols, alcohols, esters, ethers, polyethers, carbonates, ketones, nitriles, amides, carbamates, sulphoxides, and salts of amines and quaternary ammonium compounds and a second compound selected from the group consisting of alkanols having from 6 to 18 carbon atoms, an alkyl phenol in which the alkyl group contains from 7 to 12 carbon atoms, and tributylphosphate.

The aforementioned additional modifiers may be used in the preparation of extractant compositions containing one or more orthohydroxyarylaloximes or orthohydroxyarylketoimes, one or more esters substituted with a hydroxyl group, one or more additional modifiers and a water immiscible organic solvent.

According to a second aspect of the present invention, there is provided a process for the extraction of a metal from solution in which an acidic solution containing a dissolved metal is contacted with a solvent extraction composition comprising a water immiscible organic solvent and a water-immiscible solvent extractant, whereby at least a fraction of the metal is extracted into the organic solution, characterised in that the solvent extraction composition comprises one or more orthohydroxyarylaloximes or orthohydroxyarylketoimes and one or more esters substituted with a hydroxyl group.

Metals that may be extracted in the process according to the second aspect of the present invention include copper, cobalt, nickel, manganese and zinc, most preferably copper.

The orthohydroxyarylaldoxime, orthohydroxyarylketoxime, esters substituted with a hydroxyl group, and water immiscible organic solvent are as herein described before. It is preferred that there is predominance of orthohydroxyarylaldoximes in relation to any orthohydroxyarylketoximes present in the solvent extraction composition. It is especially preferred that the solvent extraction composition comprises one or more orthohydroxyarylaldoximes and one or more esters substituted with a hydroxyl group.

The aqueous acidic solution from which metals are extracted by the process of the second aspect of the present invention often has a pH in the range of from -1 to 7, preferably from 0 to 5, and most preferably from 0.25 to 3.5. Preferably, when the metal to be extracted is copper pH values of less than 3 chosen so that the copper is extracted essentially free of iron, cobalt or nickel. The solution can be derived from the leaching of ores or may be obtained from other sources, for example metal containing waste streams such as from copper etching baths.

The concentration of metal, particularly copper, in the aqueous acidic solution will vary widely depending for example on the source of the solution. Where the solution is derived from the leaching of ores, the metal concentration is often up to 75g/l and most often from 1 to 40g/l. Where the solution is a waste stream, the metal concentrations are often somewhat higher than those from the leaching of ores, for example up to 150g/l, usually from 75 to 130g/l.

Preferred solvent extraction compositions are those aldoxime containing compositions as discussed above with respect to the first aspect of the present invention.

The process of the second aspect of the present invention can be carried out by contacting the solvent extractant composition with the aqueous acidic solution. Ambient or elevated temperatures, such as up to 75°C can be employed if desired. Often a temperature in the range of from 5 to 60°C, and preferably from 15 to 40°C, is employed. The aqueous solution and the solvent extractant are usually agitated together to maximise the interfacial areas between the two solutions. The volume ratio of solvent extractant to aqueous solution are commonly in the range of from 20:1 to 1:20, and preferably in the range of from 5:1 to 1:5. In many embodiments, to reduce plant size and to maximise the use of solvent extractant, organic to aqueous volume ratios close to 1:1 are employed, such as 1.5:1 or less.

The mole ratio of the total of orthohydroxyarylaldoxime and/or orthohydroxyarylketoxime present to copper transferred is often selected to be in the range of from 2.7:1 to 1:1. Preferably, to achieve improved hydrometallurgical properties, such as reduced viscosity and improved phase disengagement, the mole ratio of oxime to copper transferred is from 2.3:1 to 2.0:1.

After contact with the aqueous acidic solution, the metal can be recovered from the solvent extractant by contact with an aqueous acidic strip solution.

The aqueous strip solution employed in the process according to the second aspect of the present invention is usually acidic, commonly having a pH of 2 or less, and preferably a pH of 1 or less, for example, a pH in the range of from -1 to 0.5. The strip solution commonly comprises a mineral acid, particularly sulphuric acid, nitric acid or hydrochloric acid. In many embodiments, acid concentrations, particularly for sulphuric acid, in the range of from 130 to 200g/l and preferably from 150 to 180g/l are employed. A low acid concentration but at least 4M chloride containing strip solution as described in European Patent application no. 93301095.1 (publication no. 0 562 709 A2) or International application publication No. WO95/04835 (both of which are incorporated herein by reference) can be employed. When the extracted metal is copper or zinc, preferred strip solutions respectively comprise stripped or spent electrolyte from a copper or zinc electro-winning cell, typically comprising up to 80g/l copper or zinc, often greater than 20g/l copper or zinc and preferably from 30 to 70g/l copper or zinc, and up to 220g/l sulphuric acid, often greater than 120g/l sulphuric acid, and preferably from 150 to 180g/l sulphuric acid.

The volume ratio of organic solution to aqueous strip solution in the process of the second aspect of the present invention is commonly selected to be such so as to achieve transfer, per litre of strip solution, of up to 50g/l of metal, especially copper into the strip solution from the organic solution. In many industrial copper electrowinning processes transfer is often from 10g/l to 35g/l, and preferably from 15 to 20g/l of copper per litre of strip solution is transferred from the organic solution. Volume ratios of organic solution to aqueous solution of from 1:2 to 15:1 and preferably from 1:1 to 10:1, especially less than 3:1 are commonly employed.

A preferred embodiment of the second aspect of the present invention comprises a process for the extraction of a metal from aqueous acidic solution in which:

in step 1, a water-immiscible solvent extraction composition comprising an orthohydroxyarylaloxime and one or more esters substituted with a hydroxyl group is first contacted with the aqueous acidic solution containing metal,

in step 2, separating the solvent extraction composition containing metal-solvent extractant complex from the aqueous acidic solution;

in step 3, contacting the solvent extraction composition containing metal-solvent extractant complex with an aqueous acidic strip solution to effect the stripping of the copper from the water immiscible phase;

in step 4, separating the metal-depleted solvent extraction composition from the loaded aqueous strip solution.

According to a third aspect of the present invention, there is provided a process for the extraction of a metal from solution in which an aqueous ammoniacal solution

containing a dissolved metal is contacted with a solvent extraction composition comprising a water immiscible organic solvent and a water-immiscible solvent extractant, whereby at least a fraction of the metal is extracted into the organic solution, characterised in that the solvent extraction composition comprises one or more orthohydroxyarylaloximes or orthohydroxyarylketoimes and one or more esters substituted with a hydroxyl group.

Metals that may be extracted in the process according to the third aspect of the present invention include copper, cobalt, nickel, manganese and zinc, most preferably copper.

The orthohydroxyarylaloxime, orthohydroxyarylketoime, esters substituted with a hydroxyl group and water immiscible organic solvent are as herein described before. It is preferred that there is predominance of orthohydroxyarylketoimes in relation to any orthohydroxyarylaloximes present in the solvent extraction composition. It is especially preferred that the solvent extraction composition comprises one or more orthohydroxyarylketoime and one or more esters substituted with a hydroxyl group.

The aqueous ammoniacal solution from which metals are extracted by the process of the third aspect of the present invention often has a pH in the range of from 7 to 12, preferably from 8 to 11, and most preferably from 9 to 10. The solution can be derived from the leaching of ores, particularly chalcocite ores, or may be obtained from other sources, for example metal containing waste streams such as from copper etching baths.

Preferred solvent extraction compositions are those ketoime containing compositions as discussed above with respect to the first aspect of the present invention.

The concentration of metal, particularly copper, in the aqueous ammoniacal solution will vary widely depending for example on the source of the solution. Where the solution is derived from the leaching of ores, the metal concentration is often up to 75g/l and most often from 10 to 40g/l. Where the solution is a waste stream, the metal concentrations are often somewhat higher than those from the leaching of ores, for example up to 150g/l, usually from 75 to 130g/l.

The process of the third aspect of the present invention can be carried out by contacting the solvent extractant composition with the aqueous ammoniacal solution. Ambient or elevated temperatures, such as up to 75°C can be employed if desired. Often a temperature in the range of from 15 to 60°C, and preferably from 30 to 50°C, is employed. The aqueous solution and the solvent extractant are usually agitated together to maximise the interfacial areas between the two solutions. The volume ratio of solvent extractant to aqueous solution are commonly in the range of from 20:1 to 1:20, and preferably in the range of from 5:1 to 1:5. In many embodiments, to reduce plant size and to maximise the use of solvent extractant, organic to aqueous volume ratios close to 1:1 are employed, such as 1.5:1 or less, and preferably 1.3:1 or less.

The mole ratio of the combined orthohydroxyarylaloxime and orthohydroxyarylketoxime present to copper transferred is often selected to be in the range of from 2.7:1 to 1:1. Preferably, to achieve improved hydrometallurgical properties, such as reduced viscosity and improved phase disengagement, the mole ratio of oxime to copper transferred is from 2.3:1 to 2.0:1.

After contact with the aqueous ammoniacal solution, the metal can be recovered from the solvent extractant by contact with an aqueous strip solution having a pH lower than that from which the metal was extracted.

The aqueous lower pH strip solution employed in the process according to the third aspect of the present invention is usually acidic and is as described for the strip solution in the process of the second aspect of the present invention. When the extracted metal is copper or zinc, preferred strip solutions respectively comprise stripped or spent electrolyte from a copper or zinc electro-winning cell, typically comprising up to 80g/l copper or zinc, often greater than 40g/l copper or zinc and preferably from 50 to 70g/l copper or zinc, and up to 220g/l sulphuric acid, often greater than 120g/l sulphuric acid, and preferably from 150 to 180g/l sulphuric acid.

The volume ratio of organic solution to aqueous strip solution in the process of the third aspect of the present invention is commonly selected to be such so as to achieve transfer, per litre of strip solution, of up to 50g/l of metal, especially copper into the strip solution from the organic solution. In many industrial copper electrowinning processes transfer is often from 10g/l to 35g/l, and preferably from 15 to 20g/l of copper per litre of strip solution is transferred from the organic solution. Volume ratios of organic solution to aqueous solution of from 1:2 to 15:1 and preferably from 1:1 to 10:1, especially less than 3:1 are commonly employed.

A preferred embodiment of the third aspect of the present invention comprises a process for the extraction of a metal from aqueous ammoniacal solution in which:

in step 1, a water-immiscible solvent extraction composition comprising an orthohydroxyarylketoxime and one or more esters substituted with a hydroxyl group is first contacted with the aqueous ammoniacal solution containing metal,

in step 2, separating the solvent extraction composition containing metal-solvent extractant complex from the aqueous ammoniacal solution;

in step 3, contacting the solvent extraction composition containing metal-solvent extractant complex with an aqueous strip solution of lower pH than the ammoniacal solution to effect the stripping of the copper from the water immiscible phase;

in step 4, separating the metal-depleted solvent extraction composition from the lower pH aqueous solution.

The metal can be recovered from the aqueous strip solution by conventional methods, for example by electrowinning.

The invention is further illustrated, but not limited, by the following examples.

Example General Methods

Each modifier was tested with the same oxime to determine the amount of modifier required to achieve a target minimum strip level under given strip conditions. These target levels were chosen based on experience of the performance of commercial extractants under such conditions. For the three aqueous strip solutions (compositions in parenthesis) the target minimum strip levels were 1.8gpl Cu (30gpl Cu/120gpl H_2SO_4), 1.6gpl Cu (30gpl Cu/150gpl H_2SO_4) and 1.24gpl Cu (30gpl Cu/180gpl H_2SO_4). Then using the concentrations of modifier defined by the target minimum strip level experiment, a maximum load value was determined under fixed loading conditions. Then finally by performing a strip experiment on the maximum load solutions, and calculating the difference between the loaded and stripped copper values, a net copper transfer value was derived for each modifier under fixed extract and strip conditions.

Modifier Concentration required to achieve Target Minimum Strip Value:

Sample Solutions:

5-Nonyl-2-hydroxybenzaldoxime (hereinafter "oxime") (100g) was dissolved in the hydrocarbon solvent commercially available under the trade name ORFOM SX7 (1l) to give a 100g/l stock solution. Then for each modifier to be tested, four separate aliquots (25ml) of the stock solution were removed and these were used to prepare the test solutions as follows:-

- 1) To the first aliquot (25ml) of stock solution was added 2.5g modifier and the resulting solution was then diluted to 50ml by the addition of ORFOM SX7. This gave a 50g/l oxime, 50g/l modifier test solution.
- 2) To the second aliquot (25ml) of stock solution was added 1.5g modifier and the resulting solution was then diluted to 50ml by the addition of ORFOM SX7. This gave a 50g/l oxime, 30g/l modifier test solution.
- 3) To the third aliquot (25ml) of stock solution was added 0.75g modifier and the resulting solution was then diluted to 50ml by the addition of ORFOM SX7. This gave a 50g/l oxime, 15g/l modifier test solution.
- 4) To the first aliquot (25ml) of stock solution no modifier was added, the solution was only diluted to 50ml by the addition of ORFOM SX7. This gave a 50g/l oxime, no modifier test solution.

Evaluation procedure:

For each of the modifiers, the four test solutions were each evaluated by stripping with three copper/acid strip solutions. For this procedure, three aliquots (10ml) of each test solution were removed separately. Each aliquot was then loaded to equilibrium with

Cu²⁺ by shaking the test aliquot with an equal volume (1:1 O/A (organic/aqueous)) of an aqueous CuSO₄ solution comprising 10g/l Cu²⁺ buffered at pH4.5 for two minutes in a separating funnel, discarding the aqueous phase and repeating with fresh copper solution three further times. Once loaded with copper, each organic phase was isolated and then stripped of copper. Stripping of each solution was carried out by shaking the test aliquot with an equal volume of a given strip solution for two minutes in a separating funnel, discarding the aqueous phase and repeating with fresh strip solution three further times. Aqueous strip solutions consisting of 30gpl Cu/120gpl H₂SO₄, 30gpl Cu/150gpl H₂SO₄ and 30gpl Cu/180gpl H₂SO₄ were used for these evaluations. The copper concentration of the resulting organic phases were determined by atomic absorption spectroscopy. In total twelve measurements were made for each modifier tested.

The data gathered for each of the strip solutions employed was plotted to give a curve showing copper left in the organic phase post stripping versus concentration of modifier required. This was then used to determine the required amount of each modifier to achieve the target minimum strip level under each of the stripping conditions.

Net Copper Transfer:

Test Solutions:

Three test solutions were prepared for each modifier containing 50g/l oxime and modifier at the concentration determined to achieve the target minimum strip values. These were prepared from aliquots (25ml) of stock solution (nonylsalicylaldoxime (100g) dissolved in ORFOM SX7 (1l) to give a 100g/l stock solution) and by adding the required amount of modifier then diluting to 50ml by the addition of ORFOM SX7.

Evaluation Procedure:

Maximum Load Value:

Aliquots (10ml) of each test solution were removed separately and each loaded to equilibrium with copper by contacting with an aqueous CuSO₄ solution comprising 5g/l Cu²⁺ at 1:1 O/A by shaking 4 times for 2min in a separating funnel with fresh loading solution. Copper loading solutions of pH 1.0, 1.5 and 2.0 were used in the evaluation. Five loaded organic solutions were prepared for each modifier, one at pH 1.0, one at pH 1.5 and three at pH 2.0. The organic layers were separated and filtered through phase separation media before being analysed for copper content by atomic absorption spectroscopy.

Strip Value:

The three organic solutions for each modifier loaded at pH 2.0 were then stripped, a different strip condition being used for each. Stripping was carried out at 1:1 O/A by

contacting an aliquot (5ml) of the loaded organic solution with aqueous strip solutions comprising 30gpl Cu/120gpl H₂SO₄, 30gpl Cu/150gpl H₂SO₄ or 30gpl Cu/180gpl H₂SO₄ (contact was by shaking 4 times for 2min in a separating funnel with fresh strip solution). Copper stripping was determined by measuring copper levels in the organic phase by atomic absorption spectroscopy.

As the minimum stripping that can be achieved by the strip conditions is independent of the load conditions, the Net Copper Transfer for a combination of load and strip conditions was determined for each modifier/oxime composition by subtracting the Strip Value for a given strip solution from the Load Value at a given pH.

Example 1

The following results were obtained for the Hydroxy-ester, 2,2,4-trimethyl-1,3-pentanediol mono-isobutyrate.

The levels of modifier required to achieve the target minimum strip values were 27gpl (30/180), 33gpl (30/150) and 38gpl (30/120).

Using these modifier concentrations a matrix of net copper transfer values under the given pH loading conditions and the strip conditions were generated and are shown in Table 1.

gpl modifier	pH1 - 30/120	pH1- 30/150	pH1- 30/180	pH1.5 - 30/120	pH1.5 - 30/150	pH1.5 - 30/180	pH2 - 30/120	pH2 - 30/150	pH2 - 30/180
27	3.09	3.54	3.96	3.68	4.19	4.55	3.83	4.28	4.7
33	3.33	3.72	4.1	3.99	4.38	4.765	4.07	4.46	4.845
38	3.405	3.775	4.11	4.155	4.525	4.86	4.325	4.695	5.03

Table 1

Comparative Example A

The following results were obtained for a known commercial modifier 2,2,4-trimethyl-1,3-pentanediol di-isobutyrate.

The levels of modifier required to achieve the target minimum strip values were 41gpl (30/180), 48gpl (30/150) and 55gpl (30/120).

Using these modifier concentrations a matrix of net copper transfer values under the given pH loading conditions and the strip conditions were generated and are shown in Table 2.

gpl modifier	pH1 - 30/120	pH1- 30/150	pH1- 30/180	pH1.5 - 30/120	pH1.5 - 30/150	pH1.5 - 30/180	pH2 - 30/120	pH2 - 30/150	pH2 - 30/180
41	2.71	3.26	3.66	3.53	4.08	4.48	3.65	4.2	4.6
48	2.925	3.425	3.725	3.675	4.17	4.475	3.85	4.35	4.65
55	3.08	3.58	3.88	3.85	4.35	4.65	4	4.5	4.8

Table 2

The results show that less 2,2,4-trimethyl-1,3-pentanediol mono-isobutyrate (a
 5 modifier according to the present invention) is required to achieve the minimum strip
 value. Furthermore, a point by point comparison of the data presented for each modifier,
 shows that under each set of load/strip conditions, not only is less Hydroxy-ester modifier
 required, a higher Net Copper Transfer is achieved and hence the composition of the
 present invention is a more effective copper transfer agent than the comparative
 10 commercial composition.

CLAIMS

1. A solvent extraction composition comprising one or more orthohydroxyarylaloximes or orthohydroxyarylketoximes and one or more esters substituted with a hydroxy group.

2. A composition according to claim 1, additionally comprising a water immiscible organic solvent.

3. A composition according to either preceding claim, wherein the orthohydroxyaryl oxime or orthohydroxyarylketoxime is selected from the class of compounds represented by the Formula (1),



Formula (1)

wherein

R¹ is hydrogen or an optionally substituted hydrocarbyl group

R² is an optionally substituted ortho-hydroxyaryl group, and salts thereof.

4. A composition according to any one of claims 1 to 3, wherein the ester substituted with a hydroxy group is selected from the class of compounds represented by formula:



wherein one of R⁷ or R⁸ is a substituted hydrocarbyl group with at least one hydroxyl group and the other is an optionally substituted hydrocarbyl group.

5. A composition according to any one of claims 1 to 4, wherein the orthohydroxyarylketoxime is a 5-(C₉ to C₁₄ alkyl)-2-hydroxyacetophenone oxime, preferably 5-nonyl-2-hydroxy-acetophenone oxime.

6. A composition according to any one of claims 1 to 5, wherein the orthohydroxyarylaloxime is a 5-(C₉ to C₁₄ alkyl)-2-hydroxybenzaloxime, preferably 5-nonyl-2-hydroxy-benzaloxime.

7. A composition according to any one of claims 1 to 6, wherein the ester substituted with a hydroxy group comprises a highly-branched hydroxy-ester comprising from 9 to 25 carbon atoms.

8. A composition according to any one of claims 1 to 7, wherein the hydroxy functionality of the ester substituted with a hydroxy group resides on R⁸, and where R⁸ is branched aliphatic group.

9. A composition according to any preceding claim wherein the ester substituted with a hydroxy group is 2,2,4-trimethyl-1,3-pentanediol mono-isobutyrate or 2,2,4-trimethyl-1,3-pentanediol monobenzoate

10. A process for the extraction of a metal from solution in which either an acidic solution containing a dissolved metal or an aqueous ammoniacal solution containing a dissolved metal is contacted with a solvent extraction composition comprising a water immiscible organic solvent and a water-immiscible solvent extractant, whereby at least a fraction of the metal is extracted into the organic solution, characterised in that the solvent extraction composition comprises one or more orthohydroxyarylaloximes or orthohydroxyarylketoimes and one or more esters substituted with a hydroxy group.

11. A process according to claim 10 wherein there is a predominance of orthohydroxyarylaloximes in relation to any orthohydroxyarylketoimes present in the solvent extraction composition

12. A process according to claim 10 wherein there is a predominance of orthohydroxyarylketoimes in relation to any orthohydroxyarylaloximes present in the solvent extraction composition.

13. A process according to any one of claims 10 to 12, wherein the metal is copper, zinc, cobalt or nickel, and is preferably copper.

14. A process according to any one of claims 10 to 13, wherein the orthohydroxyaryl oxime or orthohydroxyarylketoime is selected from the class of compounds represented by the Formula (1),



Formula (1)

wherein

R¹ is hydrogen or an optionally substituted hydrocarbyl group

R² is an optionally substituted ortho-hydroxyaryl group,

and salts thereof.

15. A process according to any one of claims 10 to 14, wherein the ester substituted with a hydroxy group is selected from the class of compounds represented by formula:



Formula (2)

5 wherein one of R^7 or R^8 is a substituted hydrocarbyl group with at least one hydroxyl group and the other is an optionally substituted hydrocarbyl group

16. A process according to any one of claims 10 to 15, wherein the orthohydroxyarylketo xime is a 5-(C_6 to C_{14} alkyl)-2-hydroxyacetophenone oxime, preferably 5-nonyl-2-hydroxy-acetophenone oxime.

17. A process according to any one of claims 10 to 16, wherein the orthohydroxyaryldoxime is a 5-(C_6 to C_{14} alkyl)-2-hydroxybenzaldoxime, preferably 5-nonyl-2-hydroxy-benzaldoxime.

18. A process according to any one of claims 10 to 17, wherein the ester substituted with a hydroxy group comprises a highly-branched hydroxy-ester comprising from 9 to 25 carbon atoms.

19. A process according to any one of claims 10 to 18, wherein the hydroxy functionality of the ester substituted with a hydroxy group resides on R^8 , and where R^8 is a branched aliphatic group.

20. A process according to any one of claims 10 to 19, wherein the ester substituted with a hydroxy group is 2,2,4-trimethyl-1,3-pentanediol mono-isobutyrate or 2,2,4-trimethyl-1,3-pentanediol monobenzoate.

ABSTRACTCOMPOSITION AND PROCESS FOR THE EXTRACTION OF METALS

A solvent extraction composition is provided which comprises one or more orthohydroxyarylaloximes or orthohydroxyarylketoximes and one or more esters substituted with a hydroxyl group, and preferably a water immiscible organic solvent. The orthohydroxyarylaloximes, or orthohydroxyarylketoximes commonly have Formula (1),



Formula (1)

wherein R^1 is hydrogen or a hydrocarbyl group, and R^2 is an ortho-hydroxyaryl group; and the esters substituted with a hydroxyl group are of Formula (2),



Formula (2)

wherein one of R^7 or R^8 is a substituted hydrocarbyl group with at least one hydroxyl group and the other is an optionally substituted hydrocarbyl group. Preferred orthohydroxyarylaloximes are 5-(C_9 to C_{14} alkyl)-2-hydroxybenzaloximes and preferred orthohydroxyarylketoximes are 5-(C_9 to C_{14} alkyl)-2-hydroxyacetophenone oximes. Preferred esters substituted with a hydroxy group are highly-branched alkyl esters comprising from 5 to 51 carbon atoms, wherein the hydroxy group resides on R^8 . Processes for the extraction of metal values from aqueous acidic and ammoniacal solutions are also provided.

RULE 63 (37 C.F.R. 1.63)
DECLARATION AND POWER OF ATTORNEY FOR PATENT APPLICATION IN THE
UNITED STATES PATENT AND TRADEMARK OFFICE

As a below named inventor, I hereby declare that my residence, post office address and citizenship are as stated below next to my name and I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the INVENTION ENTITLED:

Composition of oxime and hydroxy-ester for the solvent extraction of metals

the specification of which

is attached hereto
 was filed on as U.S. application serial No.
 X was filed as PCT international application No. PCT/GB99/03807 on 16/11/1999
 and (if applicable) was amended on

I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above. I acknowledge the duty to disclose all information which is known to me to be material to patentability as defined in 37 C.F.R. 1.56. I hereby claim foreign priority benefits under 35 U.S.C. 119/365 of any foreign application(s) for patent or inventor's certificate listed below and have also identified below any foreign application for patent or inventor's certificate filed by me or my assignee disclosing the subject matter claimed in this application and having a filing date (1) before that of the application on which priority is claimed, or (2) if no priority is claimed, before the filing date of this application:

PRIOR FOREIGN APPLICATION(S)

<u>Number</u>	<u>Country</u>	<u>Day/MONTH/Year Filed</u>	<u>Date First Laid Open or published</u>	<u>Date Patented or Granted</u>	<u>Priority claimed</u>	
					<u>Yes</u>	<u>No</u>
9827288.3	United Kingdom	12/12/1998				X

I hereby claim the benefit under 35 U.S.C. 120/365 of all United States applications listed below and PCT international applications listed above or below and, if this is a continuation-in-part (CIP) application insofar as the subject matter disclosed and claimed in this application is in addition to that disclosed such in the prior applications. I acknowledge the duty to disclose all information known to

me to be material to patentability as defined in 37 C.F.R. 1.56 which became available between the filing date of each such prior application and the national or PCT international filing date of this application:

PRIOR U.S. OR PCT APPLICATION(S)

Application No. (Serial Code/Serial No.) Day/MONTH/Year Filed

Status
(patented, pending abandoned)

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

And I hereby appoint Cushman, Darby & Cushman, L.L.P. 1100 New York Avenue, N.W. Ninth Floor, East Tower, Washington, D.C. 20005-3918, telephone number 861-3000 (to whom all communications should be directed), and the below named persons (of the same address) individually and collectively my attorneys to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith and with the resulting patent and I hereby authorize them to act and reply on instructions from and communicate directly with the person/assignee/attorney/firm/organisation who/which first sends/sent this case to them and by who/which I hereby declare that I have consented after full disclosure to be represented unless/until I instruct Cushman, Darby and Cushman in writing to the contrary.

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